

General Stats

Tophat

Bowtie 2 / HiSAT2



A modular tool to aggregate results from bioinformatics analyses across many samples into a single report.

Report generated on 2022-04-23, 10:40 based on data in: /home/khoidnyds/RNAseq_old/6.tophat2

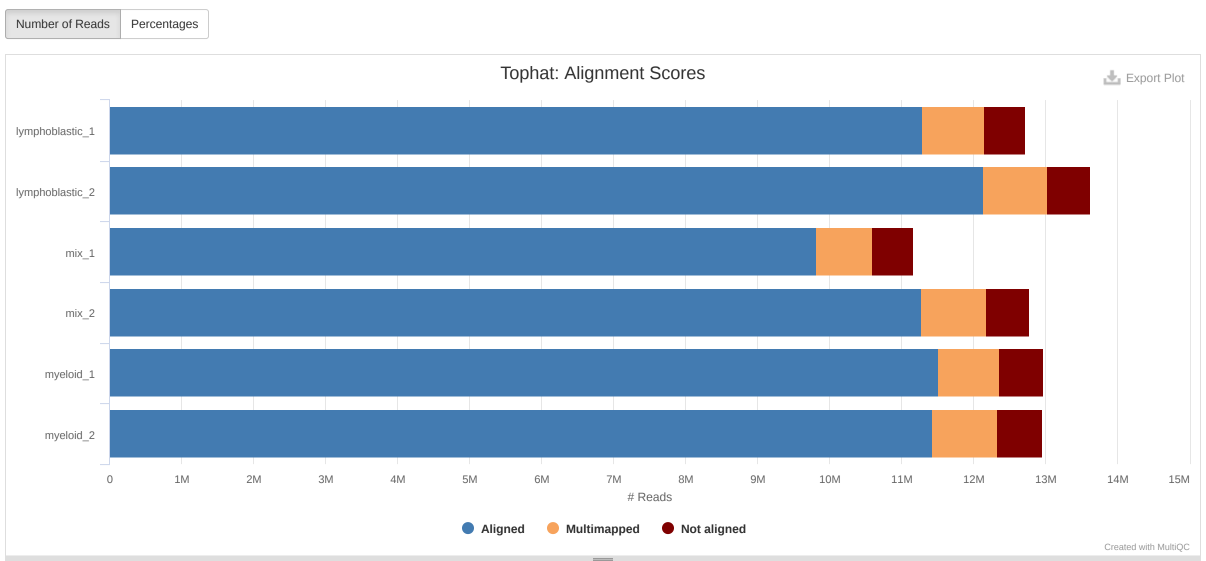
3 Welcome! Not sure where to start? Watch a tutorial video (6:06) don't show again ★

General Statistics

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|---|-----------|-----------|-----------|
| ☐ Copy table ☐ Configure Columns ☐ Plot ☐ Showing 12/12 rows and 3/3 columns. | | | |
| Sample Name | % Aligned | M Aligned | % Aligned |
| bowtie.left_kept_reads | | | 83.9% |
| bowtie.left_kept_reads_seg1 | | | 35.0% |
| bowtie.left_kept_reads_seg2 | | | 50.4% |
| bowtie.left_kept_reads_seg3 | | | 54.1% |
| bowtie.left_kept_reads_seg4 | | | 53.0% |
| bowtie.left_kept_reads_seg5 | | | 29.1% |
| lymphoblastic_1 | 95.5% | 11.3 | |
| lymphoblastic_2 | 95.6% | 12.1 | |
| mix_1 | 94.9% | 9.8 | |
| mix_2 | 95.3% | 11.3 | |
| myeloid_1 | 95.2% | 11.5 | |
| myeloid_2 | 95.1% | 11.4 | |

Tophat

 $To phat is a fast splice junction mapper for RNA-Seq reads. It aligns RNA-Seq reads to mammalian-sized genomes. \ \textit{DOI: 10.1186/gb-2013-14-4-r36; 10.1093/bioinformatics/btp120.} \\$



Bowtie 2 / HiSAT2

Bowtie 2 and HISAT2 are fast and memory-efficient tools for aligning sequencing reads against a reference genome. Unfortunately both tools have identical log output by default, so it is impossible to distiguish which tool was used. . DOI: 10.1038/nmeth.1923; 10.1038/nmeth.3317; 10.1038/s41587-019-0201-4.



